# WIEN

## Indium Mediated Acyloxyallylation of Aldoses

Lara Maltrovsky, Nina Biedermann, Markus Draskovits, Christian Stanetty\*

Institute of Applied Synthetic Chemistry, TU Wien, Getreidemarkt 9,1060 Vienna, Austria

#### State of the Art: Indium Mediated Acyloxyallylation (IMA) of (Un)protected Aldoses

The indium mediated acyloxyallylation of aldoses was introduced by Madsen et al. in 2005 and constitutes a useful tool for the elongation of reducing sugars by two carbon atoms (upon ozonolysis).<sup>1</sup> While the studied transformation can lead to four different diastereomers, the mainly observed isomer was the product with lyxo-configuration, representing a syn-orientation in respect to the former 2-OH-group and an anti-addition to the aldehyde. The latter had been expected<sup>2</sup>, however, the predominant syn-orientation in regard to the former C2-stereocenter is a specific feature for reducing sugars as starting materials and was exploited in our case study towards bacterial L-glycero-D-manno heptose at scale.<sup>3</sup> Following up, we engaged in an in-depth methodological investigation of unprotected L-erythrose and a protected version thereof which furnished a pronounced diastereodivergence.<sup>4</sup> This study is now expanded to the more generally accessible class of protected sugar aldehydes.

lyxo = *syn/anti* 



#### **Products of the IMA**

Due to the introduction of two new stereocenters during the indium mediated elongation reaction four different diastereomers can be formed in total:



#### **IMA of Protected Sugar Aldehydes**

The goal of the project was the investigation of the diastereomeric ratio of the formed products in respect to the used protecting group. The diastereomeric ratios were determined from the <sup>1</sup>H-NMR spectra of the obtained product mixtures upon complete deprotection.



protecting	<b>A</b>	<b>B</b>	<b>C</b>	<b>D</b>
group	syn/anti	syn/syn	antilanti	anti/syn
-	58%	35%	7%	

### **Synthesis of Protected Sugar Aldehydes**

We chose protected sugar aldehydes as substrates as they can be synthesized from all sugars alike and selected D-ribose as stereochemical pattern. The different derivatives were afterwards subjected to the indium mediated acyloxyallylation to investigate the influence of a variety of commonly used protecting groups onto the diastereoselectivity.



3,4- acetonide	60%	24%	10%	6%
2,3; 4,5- acetonide	14%	16%	53%	17%
acetate	14%	15%	32%	39%
benzoate	17%	15%	30%	38%
TBS	22%	22%	27%	29%

Compound A with *lyxo*-configuration (*syn/anti*) was found to be the main isomer for D-ribose and also 3,4-O-isopropylidene-D-ribose, probably due to the chelation with the OH-group on C2. In contrary, full protection of the reducing sugar led to a shift towards formation of enitols **C** and **D** which show *ribo*- (*anti/anti*) and *arabino*-configuration (*anti/* syn), respectively. However, of the used protecting groups only the acetonide protection delivers a reasonable opposing selectivity of the elongation reaction towards a major product with *ribo*-configuration.

#### **Conclusion and Outlook**

We have successfully prepared a family of *ribo*-configured protected sugar aldehydes and subjected them to the indium mediated acyloxyallylation. Within our investigation we found diastereodivergence depending on whether the starting materials (2-OH) did bear protecting groups or not. The effect of different protecting groups was examined and a protection via acetonides was found to be most suitable to shift the reaction outcome towards an *anti/anti*-configured product.

TBSO OTBS

BzO OBz

#### Indium Mediated Acyloxyallylation

Reagents and conditions: (a) NH<sub>2</sub>OMe<sup>·</sup>HCl, pyridine, rt; (b) Ac<sub>2</sub>O, DMAP, pyridine, rt; (c) BzCl, pyridine, rt; (d) O<sub>3</sub>, DCM, DMS, -80 °C to rt; (e) C<sub>3</sub>H<sub>8</sub>S, conc. HCl, 0 °C to rt; (f) DMP, *p*-TsOH, acetone, rt; (g) TBDMS triflate, 2,6-lutidine, dry DCM, 0 °C to rt; (h) I<sub>2</sub>, NaHCO<sub>3</sub>, acetone/water, rt; (i) DMP, Amberlyst® 15(H), DMF, rt.

Our study will be expanded to at least one starting material with *threo*'-configuration to confirm the generality of our findings and the effect of the most promising acetonide protection will be attempted to be boosted by larger acetal protecting groups. The successful application of the reaction on protected aldoses is encouraging and we hope that our findings help to promote a more frequent consideration of the indium mediated acyloxyallylation as a suitable synthetic solution for a two-carbon elongation within and even beyond the scope of carbohydrate chemistry.

1. Palmelund, A.; Madsen, R., Chain Elongation of Aldoses by Indium-Mediated Coupling with 3-Bromopropenyl Esters, J. Org. Chem. 2005, 70, 8248-8251. 2. Lombardo, M.; Morganti, S.; Trombini, C., 3-Bromopropenyl Esters in Organic Synthesis: Indium- and Zinc-Mediated Entries to Alk-1-ene-3,4-diols, J. Org. Chem. 2003, 68, 997-1006. 3. Stanetty, C.; Baxendale, I., Large-Scale Synthesis of Crystalline 1,2,3,4,6,7-Hexa-O-acetyl-L-glycero-alpha-D-manno-heptopyranose, Eur. J. Org. Chem. 2015, 2718-2726. 4. Draskovits, M.; Stanetty, C.; Baxendale, I.; Mihovilovic, M., Indium- and Zinc-mediated Acyloxyallylation of Protected and Unprotected Aldotetroses, J. Org. Chem. 2018, 83, 2647-2659.